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85 **Abstract**

86 The Amyloid Precursor Protein (APP) has been extensively studied as the precursor of the β -
87 amyloid peptide (A β) peptide, the major component of the senile plaques found in the brain of
88 Alzheimer's disease (AD) patients. However, the function of APP *per se* in neuronal physiology
89 remains to be fully elucidated. APP is expressed at high levels in the brain. It resembles a cell
90 adhesion molecule or a membrane receptor, suggesting that its function relies on cell-cell
91 interaction and/or activation of intracellular signaling pathways. In this respect, the APP
92 intracellular domain (AICD) was reported to act as a transcriptional regulator. Here, we used a
93 transcriptome-based approach to identify the genes transcriptionally regulated by APP in the
94 rodent embryonic cortex and upon maturation of primary cortical neurons. Surprisingly, the
95 overall transcriptional changes were subtle, but a more detailed analysis pointed to genes
96 clustered in neuronal-activity dependent pathways. In particular, we observed a decreased
97 transcription of Neuronal PAS domain protein 4 (NPAS4) in APP^{-/-} neurons. NPAS4 is an
98 inducible transcription factor (ITF) regulated by neuronal depolarization. The down-regulation of
99 NPAS4 co-occurs with an increased production of the inhibitory neurotransmitter GABA and a
100 reduced expression of the GABA_A receptors α 1. CRISPR-Cas-mediated silencing of NPAS4
101 in neurons led to similar observations. Patch-clamp investigation did not reveal any functional
102 decrease of GABA_A receptors activity, but LTP measurement supported an increased GABA
103 component in synaptic transmission of APP^{-/-} mice. Together, NPAS4 appears to be a
104 downstream target involved in APP-dependent regulation of inhibitory synaptic transmission.

105

106 **Significance Statement**

107 The Amyloid Precursor Protein (APP) is a key player in Alzheimer's disease (AD) pathogenesis.
108 We report the down-regulation of the activity-dependent transcription factor Neuronal PAS
109 domain protein 4 (NPAS4) in APP-deficient neurons, along with an increase in GABAergic
110 neuron markers and GABA release, but not in excitatory glutamatergic markers. We identified
111 NPAS4 as an APP target gene by a transcriptome analysis of APP^{+/+} versus APP^{-/-} primary
112 cortical neurons at different stages of differentiation. The downregulation of NPAS4 observed in
113 APP^{-/-} neurons was confirmed by APP silencing with a CRISPR/Cas9 approach. CRISPR/Cas9-
114 silencing of NPAS4 mimicked APP deficiency and increased GABAergic markers. The activity-
115 dependent transcription factor NPAS4 is therefore a key downstream target in the synaptic
116 functions regulated by APP.

117

118 Introduction

119 The physiological functions of APP *per se* have been largely overlooked in comparison with its
120 role of precursor of the A β peptide. A β deposition is a central event in AD, but alterations of APP
121 physiological functions are likely to play a key role in the pathogenesis.

122 APP belongs to the APP-like protein family (with APLP1 and APLP2, referred to as APLPs),
123 expressed in most of the species. The APLP family results from several duplications and
124 contraction events during evolution. The specific functions ascribed to each member are yet not
125 clearly defined (for a review see Shariati and De Strooper, 2013). APP^{-/-} mice show a subtle
126 phenotype, with reduced body and brain weight, reduced locomotor activity, gliosis, mild axonal
127 growth/white matter defects (Guo et al., 2012; Muller et al., 2012; Muller and Zheng, 2012).
128 However, a large range of functions have been attributed to APP including neuronal proliferation,
129 differentiation (Freude et al., 2011; Hu et al., 2013) and migration during embryogenesis (Young-
130 Pearse et al., 2007). APP promotes neurite outgrowth (Hoe et al., 2009b), synapse formation
131 and activity (Priller et al., 2006; Santos et al., 2009; Lee et al., 2010; Pierrot et al., 2013;
132 Klevanski et al., 2015; Zou et al., 2016) in the central nervous system (CNS) or at the
133 neuromuscular junction (Stanga et al., 2016). APP modulates the excitatory neurotransmission
134 by interacting with AMPA (Lee et al., 2010) or NMDA receptors (Cousins et al., 2009; Hoe et al.,
135 2009a). APP deficiency reduces paired pulse depression (PPD) in mice (Seabrook et al., 1999)
136 and affects the expression of GABA receptors (Fitzjohn et al., 2000; Chen et al., 2017). Its
137 overexpression induces hyperexcitability due to failure in GABAergic neurotransmission (Born et
138 al., 2014), and triggers the GABA excitatory/inhibitory shift occurring during neuronal maturation
139 (Doshina et al., 2017).

140 Tuning inhibitory/excitatory neurotransmission is instrumental in neuronal plasticity and memory
141 formation. This process is regulated by a set of transcription factors known as inducible
142 transcription factors or ITFs. ITFs belong to the Immediate Early Genes (IEGs) family, and are

143 sensitive to neuronal activity. They control the mechanisms that “reshape” synaptic inputs on
144 neurons (West and Greenberg, 2011), and play a key role in neuronal plasticity and memory
145 formation (Alberini, 2009; Loebrich and Nedivi, 2009; Leslie and Nedivi, 2011). Among them,
146 NPAS4 is robustly expressed upon neuronal depolarization, and is involved in a transcriptional
147 program that regulates neuronal firing responses to excitatory transmission by enhancing
148 inhibition (Lin et al., 2008). Elevated activity of inhibitory neurons also induces NPAS4,
149 promoting increased excitation onto the same neurons (Spiegel et al., 2014).

150 The molecular mechanisms underlying APP functions are still elusive, but several studies
151 reported that APP function is mediated by the transcriptional regulation of so-called APP target
152 genes, which is operated by the APP intracellular domain (AICD). An increasing list of AICD
153 candidate genes has emerged from various models (reviewed in Pardossi-Piquard and Checler,
154 2012). On the other hand, APP was also reported to regulate gene transcription independently of
155 AICD release (Hicks et al., 2013; Pierrot et al., 2013). It is thus so far quite impossible to clearly
156 define (i) the precise identity of APP target genes in neurons (ii) the related molecular pathways
157 underlying APP-dependent gene transcription (iii) how APP target genes relate to APP neuronal
158 function.

159 In this study, we first aimed at identifying genes that are transcriptionally regulated by APP in
160 primary neurons. We performed a transcriptome analysis (APP^{+/+} vs. APP^{-/-}) in primary cortical
161 neurons at different stages of differentiation. Changes observed in global gene expression in the
162 absence of APP were subtle. A more detailed pathway analysis indicated that expression of
163 genes clustered in activity-dependent pathway, and among these the ITF NPAS4, were down-
164 regulated in the absence of APP after 7 days of culture. Strikingly, we observed that the amount
165 of the inhibitory neurotransmitter GABA (γ -aminobutyric acid) was increased in APP^{-/-} neurons.
166 This was supported by an increased expression of the Glutamate Decarboxylase 65 (GAD65) in
167 the same context. Glutamate levels were not altered in APP^{-/-} neurons. These observations

168 were reproduced upon acute silencing of APP by CRISPR-Cas9 editing. The knockdown of
169 NPAS4 gave similar results. Neurophysiological investigations showed that excitatory
170 postsynaptic potentials (EPSP) consecutive to a Theta-burst stimulation (TBS) decreased in
171 APP^{-/-} supporting the increase release GABA, and suggesting increased inhibitory synaptic
172 inputs in APP^{-/-} neurons. Altogether, our data provide new insight into APP-dependent neuronal
173 activity, indicating that NPAS4 is an APP downstream target gene, tuning the GABA-dependent
174 activity in neuronal networks.

175

176 **Materials and Methods**

177 ***Antibodies, chemicals and reagents***

178 All media and reagents used for cell cultures were purchased from Thermo Fisher Scientific
179 (Waltham, USA); Fetal Bovine Serum (FBS) was purchased from Biowest (Nuaillé, France).
180 Analytical grade solvents and salts were purchased from Sigma-Aldrich (St-Louis, USA). sAPP α
181 (S9564) and DAPI (D9542) were from Sigma-Aldrich (St-Louis, USA), Triton-X100 from Merck
182 (Darmstadt, Germany) and TriPure Isolation Reagent from Roche (Basel, Switzerland).
183 Microarray analysis kits were from Affymetrix (Santa Clara, USA). All reagents for RNA
184 processing or cDNA synthesis were from Bio-Rad (Hercules, USA), and primers were from
185 Sigma-Aldrich (St Louis, USA). BCA Protein Assay kit was from Thermo Fisher Scientific
186 (Waltham, USA). NuPAGE® reagents were from Invitrogen (Carlsbad, USA). PolyVinylidene
187 Fluoride (PVDF) and nitrocellulose membranes were from Merck Millipore (Billerica, USA) or
188 Amersham™ (Little Chalfont, UK). Nonfat dry milk was from Merck (Darmstadt, Germany).
189 Western Lighting® Plus-ECL reagents were from PerkinElmer (Waltham, USA) and Fluorep
190 mounting medium from bioMérieux (Marcy l'Etoile, France). Lentivirus were prepared with
191 Acrodisc® 0,45 μ m filters (Pall, NYC, USA) and LentiX™ Concentrator reagent (Clontech,
192 Mountain View, USA).

193 The following antibodies were used: APP NT 22C11 (Cat. No. MAB348, Merck Millipore,
194 Billerica, USA), anti-human APP W0-2 (Cat. No. MABN10, Merck Millipore, Billerica, USA), anti-
195 APP CT Y188 (Cat. No. ab32136 Abcam, Cambridge, UK), anti-APLP1 (Cat. No. 171615,
196 Calbiochem EMD Biosciences – Merck, Darmstadt, Germany), anti-APLP2 (Cat. No. 171616,
197 Calbiochem EMD Biosciences – Merck, Darmstadt, Germany), anti-GAPDH 14C10 (Cat. No.
198 2118, Cell Signaling, Danvers, USA), anti-MAP2 (Cat. No. M4403, Sigma-Aldrich St Louis,
199 USA), anti-GAD65 (D5G2, Cat. No. 5843 Cell Signaling, Danvers, USA), anti-mouse IgG, HRP
200 Whole antibody anti-rabbit IgG HRP (Cat. No. NA931-1ML, Amersham, Little Chalfont, UK),

201 whole goat anti-mouse antibody (Cat. No. NA934-1ML, Amersham, Little Chalfont, UK) Alexa
 202 Fluor®-488, goat anti-mouse; Alexa Fluor®-568, goat anti-rabbit; Alexa Fluor®-647 and DAPI
 203 were purchased from ThermoFisher Scientific (Waltham, USA). Glutamate assay kit was from
 204 Abcam (Cambridge, UK) and γ -aminobutyric acid (GABA) ELISA from Cloud-Clone Corporation.
 205 70 μ m Falcon™ Cell Stainers were from ThermoFisher Scientific (Waltham, USA).

206 ***Animal models***

207 All animal procedures were performed in accordance with the regulations and policies of the
 208 University animal care committee. APP^{+/+} and APP^{-/-} mice were obtained from the Jackson
 209 Laboratory (Bar Harbor, USA) as C57Bl6/J and backcrossed for > 6 generations in CD1 genetic
 210 background. Animals were housed on a 12 h light/dark cycle in standard animal care facility with
 211 access to food and water *ad libitum*. Heterozygous animals (APP^{+/-}) were bred and crossed to
 212 obtain embryos of either sex from the three different genotypes (APP^{+/+}, APP^{+/-} and APP^{-/-}) in
 213 the same litter.

214 ***Primary culture and treatments***

215 Primary cultures of cortical neurons were prepared from E18 mouse embryos. Cortices were
 216 dissected and dissociated in HBSS without calcium and magnesium and the mixture was
 217 centrifuged on FBS for 10 min at 1,000 x g to pellet cells. Cells were plated at 200,000 cells/cm²
 218 in culture dishes pre-treated with 10 μ g/ml of poly-L-lysine in phosphate buffered saline (PBS)
 219 and cultured (37°C, 5% CO₂ and humidified atmosphere). Cells were cultured for 3 to 14 days *in*
 220 *vitro* in Neurobasal® medium enriched with 2% v/v B-27® supplement medium and 1 mM L-
 221 glutamine. Half of the medium was renewed every 2-3 days. Treatments with 20 nM of soluble
 222 APP α (sAPP α) were performed for 16 h after 6 days of culture (DIV6).

223 For primary cultures of astrocytes, cortices from pups were collected at postnatal day 2 and
 224 mechanically dissociated. Astrocytes were isolated using a 30% Percoll gradient and seeded

225 into gelatin-coated tissue culture flasks. Cells were left to proliferate for 14 days at 37°C - 5%
 226 CO₂ in DMEM-glutaMAX medium supplemented with 10% FBS, 50 mg/ml penicillin–
 227 streptomycin and 50 mg/ml fungizone. Medium was renewed after 7 days, cells were passaged
 228 after 14 days and further cultured in DMEM-glutaMAX with 10% FBS. Two days after passage,
 229 FBS was reduced to 3% and medium was supplemented with the growth factor cocktail G5. All
 230 experiments/treatments were performed 7 days after, referred to as DIV7 for astrocytes. For
 231 NPAS4-induction analysis, neurons and astrocytes at DIV7 were depolarized with 50 mM KCl for
 232 2-4h.

233 ***RNA extraction, transcriptome analysis and qRT-PCR***

234 Total RNA was extracted by TriPure Isolation Reagent according to the manufacturer's protocol.
 235 RNA samples were suspended in DEPC-treated water and RNA concentration was measured
 236 (OD 260 nm) on BioSpec-nano spectrophotometer (Shimadzu Biotech). For microarray analysis,
 237 RNA quality was monitored by capillary electrophoresis using the Agilent 2100 Bioanalyzer
 238 instrument with the Agilent RNA 6000 Nano Kit (Agilent, Santa Clara, USA). 250 ng of total RNA
 239 per sample was amplified and labeled using GeneChip®WT PLUS Reagent kit (Affymetrix,
 240 Santa Clara, USA) before hybridization over night at 45°C on *GeneChip®Mouse Transcriptome*
 241 *1.0 Array*. The chip was washed on the GeneChip® Fluidics Station 450 followed by scanning
 242 on a GeneChip® Scanner on Affymetrix microarray platform. For quantitative PCR, RNA
 243 samples were reversed transcribed using iScript cDNA Synthesis Kit and real time PCR was
 244 performed in an iCycler MyIQ2 multicolor-Real-Time PCR detection system using iQ SYBR
 245 Green supermix kit (Bio-Rad, Hercules, USA). A standard curve was established for relative
 246 quantification with a fourfold dilution series (from 100 to 0.0097 ng) of a cDNA template mix.
 247 Relative quantification was calculated by the $2^{\Delta\Delta CT}$ method (*Gapdh* as housekeeping control)
 248 and then normalized (percentage or fold) to the control condition (Ct). Primer used
 249 (forward/reverse) are :

250 *Gapdh* 5'-ACCCAGAAGACTGTGGATGG-3' / 5'- ACACATTGGGGGTAGGAACA-3'

251 *Npas4* 5'-GCTATACTCAGAAGGTCCAGAAGGC-3' / 5'-TCAGAGAATGAGGGTAGCACAGC-3'

252 *Egr1* 5'-TCCTCTCCATCACATGCCTG-3' / 5'-CACTCTGACACATGCTCCAG-3'

253 *Egr3* 5'-GACTCGGTAGCCCATTACAATC-3' / 5'-ACTTTCCCAAGTAGGTCACGG-3'

254 **Western blotting**

255 Cells were solubilized and sonicated in lysis buffer (20% Glycerol, 4% SDS, 125 mM Tris-HCl
 256 pH 6.8) containing a cocktail of proteases and phosphatases inhibitors (Roche, Basel,
 257 Switzerland). When performed on tissue extracts, mice were euthanized (Ketamine/Xylazine
 258 injection) and brains were dissected after perfusion with ice cold sterile PBS. Cortices and
 259 hippocampi were isolated and quickly frozen in liquid nitrogen. Tissues were crushed using
 260 mortar pestle method. For brain protein extraction, samples were homogenized in RIPA buffer
 261 (1% (w/v) NP40, 0.5% (w/v) deoxycholic acid, 0.1% (w/v) SDS, 150 mM NaCl, 1 mM EDTA, 50
 262 mM Tris, pH 7.4) containing proteases and phosphatases inhibitors cocktail. The samples were
 263 clarified by centrifugation at 20,000 x g. Protein concentrations were determined with a BCA kit.
 264 Samples were prepared with NuPAGE LDS sample buffer (4x) and 50 mM DTT and then heated
 265 for 10 min at 70°C. 10 to 40 µg of proteins or 22 µl of culture medium were loaded per well for
 266 migration followed by transfer onto PVDF or nitrocellulose membranes. For APP C-terminal
 267 fragments, proteins were transferred on nitrocellulose (0.1 µm). Membranes were blocked in
 268 nonfat dry milk (5% in PBS, 0.05% Tween-20) and immunoblotted with anti-APP NT (22C11,
 269 1/500), anti-APP CT (Y188, 1/500), anti-APLP1 (1/1000), anti-APLP2 (1/1000) and anti-GAPDH
 270 (1/25000). Blots were revealed using ECL and signal quantification was performed using
 271 GelQuant.NET software (BiochemLabSolutions.com).

272 **ImmunoCytoFluorescence (ICF)**

Neurons grown at 100,000 cells/cm² per well on poly-L-lysine coated coverslips were fixed for 15 min in PBS/4% paraformaldehyde and washed twice in PBS for 5 min. Permeabilization and blocking steps were done in PBS/5% skimmed milk/0.3% Triton-X100. Antibodies were incubated in PBS/5% skimmed milk/0.1% Triton-X100 (M1TPBS). Primary antibodies dilutions used were: mouse anti-MAP2 (1/1000), rabbit anti-APP (Y188, 1/100) and rabbit anti-GAD65 (D5G2, 1/100). Secondary antibodies dilutions used were: goat anti-mouse Alexa Fluor®-488 (1/500), goat anti-mouse Alexa Fluor®-568 (1/500) and goat anti-rabbit Alexa Fluor®-647 (1/500). Images were acquired on Evos FL Auto microscope (Invitrogen) with GFP (Alexa Fluor®-488 or native GFP), TxRed (Alexa Fluor®-568) and CY5 (Alexa Fluor®-647) EVOS LED light cubes and analyzed with ImageJ software. For the quantification of signal area, 10X or 20X magnification images were identically thresholded for APP^{+/+} and APP^{-/-}, or Ct and CRISPR-*NPAS4*. The area of thresholded images was measured and normalized to the number of cells counted by DAPI staining. For the quantification of the APP expression intensity, image acquisition was performed using 40x objective coverslip-corrected (ThermoFischer Scientific, AMEP4699) in GFP, CY5 (APP) and DAPI channels. A total of 12, 19 and 19 images were acquired and processed to obtain 33, 46 and 51 neurons in the analysis (Figure 3B) for CRISPR control (Ct), Oligo2 and Oligo17 respectively. GFP channel images were first 8-bit transformed and thresholded to highlight only GFP staining. A region of interest (ROI) was delimited around GFP⁺ neurons in the GFP channel (green) using “wand tool” in imageJ software and transposed to CY5 (APP) channel (blue). ROI mean intensity is measured with the “Analyze” tool of ImageJ software.

AICD and CRISPR/Cas9 lentiviral constructions, production and viral transduction

Lentiviruses were used to express AICD in neurons. AICD50 tagged at the c-terminal part with hemagglutinin (HA) was cloned into pLenti CMV/TO Puro lentiviral vector (Addgene ref #17482). pLenti CMV/TO Puro empty served as a control (Ct). A lentiviral vector-based approach was

also used to deliver the CRISPR-Cas9 system. sgRNAs “Oligo2” and “Oligo17” were designed using on/off-target score algorithm to target the *APP* mouse gene (Gene ID: 11820), and sgRNA “CRISPR-*NPAS4*” to target the *NPAS4* mouse gene (Gene ID: 225872). sgRNAs were cloned in a lentiviral vector delivering sgRNA, SpCas9 and coexpressing eGFP (Addgene #57818) according to author instructions (Heckl et al., 2014). The negative Ct used was the lentiviral construct without sgRNA but expressing SpCas9 and eGFP. sgRNA used are (sequence/PAM/specificity score):

APP Oligo2 5'-GTGGAAGATCCGCCGCGCCC-3' / TGG / 95

APP Oligo17 5'-GTACCCACTGATGGCAACGC-3' / CGG / 92

Npas4 5'-GACCCTTGCGAGTGTAGATGC-3 / AGG / 83

All lentiviral vectors were validated by sequencing (Beckman Coulter Genomics, UK) prior to production and purification using the Plasmid Midi kit (Qiagen, Hilden, Germany). Production was carried out by transfecting HEK293-T cells in 10 cm dishes (2×10^6 cells/dish) with lentiviral CRISPR-Cas9 vectors, pCMV-dR8.2 (Addgene#12263) and pMD2.G (Addgene#12259). After 48 h, the supernatant was filtered and incubated with 1/3 (v/v) of LentiX™ Concentrator for 90 min on ice. The collected supernatant was centrifuged at 1,500 x g for 45 min at 4°C, the pellet was resuspended in 20 µl per dish of Neurobasal® Medium and stored at -80°C until use.

Neurons were infected with CRISPR-Cas9 lentiviruses 1 day after plating (DIV1). Typically, 20 µl of concentrated virus were used to infect 800,000 cells per well in a 12-well culture dish. The medium was completely changed after 24 hours, and a half-media change was performed every 2-3 days thereafter. The neurons were harvested at DIV7 or as indicated.

Toxicity assay

Cell viability upon lentiviral infection was measured by lactate dehydrogenase (LDH) release in the culture medium using Cytotoxicity Detection kit (Sigma-Aldrich, St-Louis, USA), according to the manufacturer's instructions. Relative absorbance was measured at 490 nm using a VICTOR

323 Multilabel Plate Reader (PerkinElmer, Richmond, USA). Background LDH release was
324 determined in non-infected control cultures.

325 ***Flow cytometry and cell sorting***

326 At DIV7, infected neurons were briefly rinsed with PBS and trypsinized for 2 min. Neurons were
327 mechanically dissociated and filtered through 70 μ m Falcon™ Cell Strainers in 50 ml tube
328 containing FBS. Cells were pelleted by centrifugation at 1,000 x g for 5 min and resuspended in
329 PBS/1% FBS/1mM EDTA. TO-PRO™-3 Iodide (Thermo Fisher Scientific, Waltham, USA) was
330 used to stain dead cells and exclude them for the sorting. Cells were sorted using a BD
331 FACSaria™III cell sorter (BD Biosciences, San Jose, USA). The sort parameters used were the
332 following: nozzle 100 μ m, sheath pressure 20 psi, drop frequency 30 kHz and sort precision 16-
333 32-0. Sample and collection tubes were maintained at 4°C throughout the procedure. GFP-
334 negative and positive cells were harvested in PBS/1% FBS/1 mM EDTA, centrifuged at 1,2000 x
335 g for 2 min and homogenized in TriPure Isolation Reagent for RNA extraction.

336 ***Glutamate and GABA measurements***

337 Neurons were grown at 200,000 cells/cm² in 12 well culture dish. Glutamate and γ -aminobutyric
338 acid (GABA) were measured in cells lysates and culture media at DIV7. Media were harvested,
339 centrifuged to pellet and remove cellular debris, treated with a cocktail of proteases inhibitors
340 and frozen at -20°C until use. Cells were scrapped in ice cold PBS, pelleted by centrifugation
341 (12,000 x g for 3 min at 4°C), quickly frozen in liquid nitrogen and kept at -80°C until use.
342 Glutamate and GABA assays were performed according to the manufacturer's protocol (Abcam,
343 Cambridge, UK). For both, cells were lysed by 5 cycles of thawing and freezing in PBS and
344 centrifuged at 12,000 x g for 10 min at 4°C. Supernatants were used for the quantification and
345 normalized on protein content. Media were directly used for quantification.

346 ***Calcium imaging***

347 Changes in intracellular $[Ca^{2+}]$ were measured in single neurons using the calcium sensitive
348 fluorescent dye Fura2 (Molecular Probes, Cambridge, UK). Neurons were grown on poly-L-
349 lysine-coated 15 mm diameter coverslips and were loaded with 2 μ M Fura2-acetoxymethyl ester
350 (Fura-2AM) for 40 min in a Krebs buffer (10 mM HEPES, 135 mM NaCl, 6 mM KCl, 2 mM
351 $CaCl_2$, 1.2 mM $MgCl_2$, 10 mM glucose, pH 7.4) at room temperature. Coverslips were rinsed
352 once and then mounted on a heated (37°C) and perfused microscope chamber (Warner
353 Instrument Corporation). While continuously perfused with heated Krebs buffer, Fura2-loaded
354 neurons were excited successively at 340 and 380 nm (excitation light was obtained from a
355 xenon lamp coupled to a monochromator) for 2x100 ms. Emitted fluorescence was monitored at
356 510 nm using a charged coupled device sensor camera coupled to an inverted Olympus IX70
357 microscope (TILL photonics). Fluorescence intensities from each single neuron was recorded
358 separately, corrected for the background and combined (fluorescence ratio F340/F380) using
359 the software TILLvisION version 3.3. Calcium signals were measured upon application
360 (perfusion) of 30 μ M glutamate (Glut) in Krebs buffer. A total of 70–80 cells were analyzed in
361 each experiment (coverslips) and non-neuronal cells were excluded from analysis as previously
362 described by Pickering and coworkers (Pickering et al., 2008). Changes in intracellular $[Ca^{2+}]$
363 were calculated from fluorescence emission intensity ratios (F340/F380). These changes were
364 expressed as normalized fluorescence where every measurement of F340/F380 was divided by
365 the basal fluorescence value corresponding to the mean of signals measured during a period of
366 25 s in basal condition (prior to the glutamate stimulation). To estimate the amplitude of the
367 response to glutamate, the area under curve (AUC) was calculated using GraphPad Prism
368 (GraphPad Software, San Diego, USA).

369 ***Field potential recordings***

370 3 months old mouse brain slices were prepared as described in (Lepannetier et al., 2018).
371 Excitatory postsynaptic potentials (EPSP) were evoked through a bipolar stimulating electrode

372 placed in the Schaffer collaterals (SC) and recorded by the AxoClamp 2B (Axon Instruments,
373 USA) amplifier through a glass electrode placed in the CA1 region (stratum radiatum). Stimuli
374 consisted of 100 μ s pulses of constant currents with intensity adjusted to produce 35% of the
375 maximum response every min. Responses were digitized by Digidata1322A (Axon Instruments,
376 USA) and recorded to a computer using WinLTP software (Anderson and Collingridge, 2007).
377 Long-term potentiation was induced by applying a theta-burst stimulation (TBS) consisting in
378 four trains of 5 pulses (100 Hz) separated by a 200 ms interval. Slopes of field excitatory
379 postsynaptic potentials (fEPSP) responses were expressed normalized to the pre-treatment
380 baseline, defined as 100%.

381 ***Electrophysiology of cultured neurons***

382 Patch-clamp recordings of primary neurons at DIV7 were carried out at room temperature, using
383 an EPC-9 amplifier controlled by PatchMaster software (HEKA Elektronik, Lambrecht,
384 Germany). GABA was applied by pressure ejection using a Picospritzer. The patch pipettes
385 were pulled with a resistance of 4–7 M Ω using a DMZ-Universal Puller (Zeitz Instruments,
386 Munich, Germany). Series resistances were compensated (75–90%) and periodically monitored.
387 The following extracellular solution was used (in mM): 140 NaCl, 5 KCl, 1 CaCl₂, 1 MgCl₂, 10
388 glucose, and 10 HEPES, pH 7.4. The pipette solution had the following composition (in mM):
389 140 CsCl, 10 EGTA, 0.3 Mg₂ATP, 0.3 CaCl₂, and 10 HEPES, pH 7.25. To prevent network
390 activity, the experiments were performed in the presence of 1 μ M tetrodotoxin (TTX), 10 μ M 6-
391 cyano-7-nitroquinoxaline-2,3-dione disodium (CNQX) and 20 μ M D-(-)-2-Amino-5-
392 phosphonopentanoic acid (D-AP5) and 100 nM CGP55845.

393 ***Statistical analysis***

394 *Microarray analysis:* Raw data were analyzed using Bioconductor (R environment). Robust
395 Multiarray Average (RMA) was used for background correction, normalization, probe level

intensity calculation and probe set summarization. Gene expression values were compared between APP^{+/+} and APP^{-/-} neurons at different stage of development DIV3, DIV7 and E18 using the R-Limma (Linear Models for MicroArray Data) package. Benjamini-Hochberg procedure was used for multiple testing corrections. Only transcripts with an Entrez ID were kept among the raw data in order to facilitate the analysis. Gene set enrichment analysis was performed on differentially expressed genes sets after the ROAST (Rotation gene set tests for complex microarray experiments) (Wu et al., 2010) procedure to identify KEGG pathways modified in absence of APP for all conditions (E18, DIV3 and DIV7). The data obtained have been deposited in NCBI's Gene Expression Omnibus (Edgar et al., 2002) and are accessible through GEO Series. Otherwise, statistical analyses were performed using GraphPad Prism (GraphPad Software, San Diego, USA). Gaussian distribution was assessed by Kolmogorov-Smirnov test (GraphPad Prism). Parametric test was applied if the data followed normal distribution. Otherwise non-parametric tests were used. When two groups were compared, parametric Student's t-test or non-parametric Mann-Whitney test were used. When more than two groups were compared, parametric ANOVA with indicated post hoc tests or non-parametric Kruskal-Wallis were used. Significance is indicated as ns = non-significant; * $p < 0.05$; ** $p < 0.01$; *** $p < 0.001$. The number of samples per condition in one experiment (n) and the number of biological replicates (N) are indicated in figure legends.

416 Results

417 APP-dependent expression of NPAS4 in differentiated primary neuron cultures

418 Transcriptome analysis were performed on primary neurons and embryonic cortex according to
419 the workflow described in Extended Data Fig. 1-1A. Neurons from embryonic cortex (E18) were
420 cultured for 3 or 7 days in vitro (DIV3 or DIV7) and up to DIV14 when necessary.
421 Characterization of APP protein family expression indicated an increase in APP, APLP1 and
422 APLP2 upon differentiation with a peak of expression at DIV7-8 (Extended Data Fig. 1-1B-C),
423 supporting an important role of APP protein family in neuronal maturation. APLP1 and APLP2
424 levels were similar in APP^{-/-} neurons and APP^{+/+} at any time point of differentiation (Extended
425 Data Fig. 1-1D). Thus, the results obtained here in APP^{-/-} neurons are related to the loss of APP
426 and not to indirect effects resulting from up- or down-regulation of APLP1 or APLP2 in the
427 absence of APP.

428 Previous studies indicated that APP-dependent gene transcription involves the release of its
429 intracellular domain or AICD. AICD is detectable in the nucleus of primary neurons at DIV6-7
430 (Kimberly et al., 2005), suggesting that AICD-dependent gene transcription should be temporally
431 restricted. We monitored AICD production at DIV7 in APP^{+/+}, APP^{+/-} and APP^{-/-} cultures
432 (Extended Data Fig. 1-1E). AICD was only readily detectable in APP^{+/+} neurons at a high
433 exposure time, confirming (i) that it is a transient peptide (Huysseune et al., 2007) with a
434 restricted temporally expression pattern in primary neurons and (ii) that APP-dependent
435 transcriptional regulation may occur at a defined time-period (around DIV7). We thus performed
436 microarray experiments at several differentiation stages (summarized in Extended Data Fig. 1-
437 1A) to track genome-wide expression changes in APP^{+/+} and APP^{-/-} primary cortical neurons
438 at DIV3 (immature neuronal network, no AICD), DIV7 (neuronal network with detectable AICD)
439 and in E18 cortical tissue (embryonic tissue). We used *Affymetrix GeneChip®Mouse*
440 *Transcriptome 1.0 Array* and performed data analysis with the R-Limma (Linear Models for

441 **MicroArray Data**) package (Ritchie et al., 2015). The chips used allowed profiling of the
 442 expression of coding and non-coding genes (lncRNA, miRNA, pseudogene...) as well as
 443 alternative splicing events. Transcriptome analysis was performed in triplicate (N=3 independent
 444 cultures) for each condition (E18, DIV3, and DIV7). We focused on differentially expressed
 445 coding genes, although data were also collected for non-coding RNAs. Strikingly, the overall
 446 changes observed (fold changes) were moderate in all conditions (E18, DIV3 and DIV7). Few
 447 coding transcripts appear to be differentially expressed when the specific fold change (linear) is
 448 set at 1.25, 1.5 or 2 (Fig. 1A). The Benjamini-Hochberg multiple correction test did not reveal
 449 any robust differential gene expression (adjusted p-value <0.05) excepted for *APP* (positive
 450 control). To note, we did not measure significant change in the expression of genes previously
 451 identified as AICD target genes (Pardossi-Piquard and Checler, 2012). Gene enrichment
 452 analysis was further performed using the ROAST (Rotation gene set test for complex microarray
 453 experiment) procedure to identify a molecular interaction/reaction networks diagram known as
 454 the KEGG pathway (Kanehisa and Goto, 2000) altered in the absence of APP. The first five
 455 pathways (in terms of significance), the number of genes modified as well as their direction are
 456 shown in Fig. 1B. For instance, *ECM (extracellular matrix)-receptor interaction* and *Long-term*
 457 *potentiation* pathways appeared to be modulated in absence of APP at DIV7. APP shares the
 458 structure of transmembrane receptors and cell adhesion proteins that activate Cell-ECM
 459 pathways. Long term potentiation (LTP) is a major mechanism in memory formation and
 460 learning. Both of these pathways have been associated to APP function (Caceres and Brandan,
 461 1997; Seabrook et al., 1999; Puzzo et al., 2011). We kept this pathways analysis to further
 462 investigate the regulation of candidate genes relevant to APP functions. In a set of arrays from a
 463 primary neuron at DIV7 (APP+/+ vs APP-/-), we noticed a down-regulation of Inducible
 464 Transcription Factors (ITFs) or Immediate Early Genes (IEGs) in APP-/- neurons (Fig. 1C).
 465 Among them, the activity-dependent transcription factor NPAS4 (Neuronal PAS domain protein
 466 4). NPAS4 is a neuron-specific ITF, known to be regulated by neuronal depolarization. We

confirmed by qPCR that the NPAS4 mRNA level was decreased at DIV7 in APP^{-/-} neurons, but neither at DIV3 nor in the cortex at E18 (Fig. 1D). To note, the expression of other IEGs (*Egr1* and *Egr3*) previously reported to be APP downstream targets (Hendrickx et al., 2013; Hendrickx et al., 2014) was not altered in our experiments (See Extended Data Fig. 1-2).

Control of NPAS4 expression by APP

Since AICD - produced at DIV7 – is reported to mediate APP nuclear signaling (Belyaev et al., 2010), we analyzed its involvement in NPAS4 expression. We transduced primary neurons with a lentiviral vector expressing the 50 C-terminal amino acids of APP (AICD) fused at the C-terminus to the hemagglutinin tag (HA). AICD-HA is detectable in infected cells (Fig. 2A) and AICD expression in APP^{-/-} neurons significantly increased NPAS4 mRNA levels (Fig. 2B), indicating that AICD is involved in the transcriptional regulation of NPAS4. As some of the APP functions were found to rely on its extracellular soluble fragment (sAPP α), we tested whether the sAPP α can regulate NPAS4 expression *per se*. Treatment of neuronal cultures with 20 nM of human sAPP α (Fig. 2C) significantly increased NPAS4 mRNA levels in APP^{+/+} neurons, but not in APP^{-/-} neurons (Fig. 2D). Together, these data indicate that (i) AICD is likely to be involved in APP-dependent NPAS4 transcription (ii) soluble APP (sAPP α) triggers NPAS4 expression, but only in a context where endogenous APP is expressed. Importantly, glial cells (about ~16% of total cells in primary cultures) could indirectly contribute to these observations. We found that the absence of APP did not change the astrocytic pattern of primary cultures, and that astrocytes did not readily express NPAS4 in contrast to neuron (Extended Data Fig. 2-1). Moreover, NPAS4 is strongly induced as expected by depolarization only in neurons (Sun and Lin, 2016). Together, this indicated that NPAS4 is a downstream transcriptional target that could be involved in APP neuronal functions.

The mild APP-dependent transcriptional regulations we observed are in line with the mild phenotype of APP knockout mice (Muller et al., 1994; Zheng et al., 1995). Still, APP-dependent

gene regulations that occur in the close-up could be hidden in the long term by functional redundancies with other members of the APP family (Shariati and De Strooper, 2013). In agreement, APP^{-/-} phenotype is better unraveled by acute down-regulation of APP in the brain (Senechal et al., 2007). We performed a knockdown of APP expression in neurons with a lentiviral-based CRISPR-Cas9 genome editing approach (Jinek et al., 2012) to test the consequence of acute APP downregulation on NPAS4 expression. Nearly ~50% of the cells were infected and no lentiviral toxicity was measured under our conditions (Extended Data Fig. 3-1). Only neuronal cells were infected, reflecting the tropism of the viral particles for neurons. APP expression was analyzed by ICF (Fig. 3A). Measurement of the intensity of APP signal in infected (GFP-positive) neurons (Fig. 3B) indicated a strong decrease in APP expression (40 to 50%) in neurons infected with CRISPR-Cas9 viruses targeting APP exon1 and exon 2 (Oligo2 and 17 sgRNA, respectively). This was confirmed by Western blotting (Fig. 3C). Importantly, expression of APLP1 and APLP2 was not altered in neurons infected under the same conditions, indicating that off target editing of APP did not occur in our experimental setup. To measure the expression of NPAS4 selectively in GFP-positive (knockdown) neurons, we sorted GFP positive neurons by flow cytometry (Fig. 3D). TO-PRO™-3 staining was used as a viability marker to exclude dead cells from the analysis. The sorting parameters were set by using non-infected neurons (GFP negative) and neurons expressing GFP (GFP positive) as standards. NPAS4 mRNA was readily decreased in neurons infected with Oligo2- and 17-expressing lentiviruses (GFP positive). Thus, acute APP knockdown resulted in decreased NPAS4 mRNA levels, confirming the APP-dependent NPAS4 transcriptional regulation observed in APP^{-/-} neurons.

APP deficiency increases markers associated to GABAergic transmission

NPAS4 is an ITF induced by neuronal activity. The down-regulation of NPAS4 expression observed in the absence of APP could reflect an impairment in the establishment of a functional network, leading to defects in basal neuronal activity. APP was reported to modulate neurite

517 outgrowth and synapse formation (Priller et al., 2006; Young-Pearse et al., 2007; Tyan et al.,
518 2012; Billnitzer et al., 2013). We analyzed neuronal arborization by measuring the area of the
519 neuron-specific microtubule associated protein2 (MAP2) signal per cell from DIV1 to DIV7 (Fig.
520 4). Neurite extension observed in APP^{+/+} and APP^{-/-} neurons was not significantly different at
521 DIV1 and DIV3. Strikingly, the absence of APP significantly increased MAP2 signal at DIV7,
522 indicating the importance of APP for in neurite arborization and formation of a functional
523 network. This observation reinforced the possible involvement of NPAS4 in APP physiological
524 function. NPAS4 activity scales the neuronal network by controlling the balance of excitatory and
525 inhibitory inputs on post-synaptic neurons (Lin et al., 2008; Bloodgood et al., 2013; Spiegel et al.,
526 2014). We measured the amount of two neuromediators, GABA (released at inhibitory
527 synapses) and glutamate (released at excitatory synapses) in the medium and in the cells of
528 primary neurons at DIV7 (Fig. 5A-B). The concentration of GABA was increased by 83% in the
529 medium of APP^{-/-} neurons (Fig. 5A). No significant change in glutamate concentration (cell or
530 medium) was observed in APP^{-/-} neuronal cultures when compared to APP^{+/+} (Fig. 5B). In line
531 with this observation, we measured only very subtle changes in glutamate responses in APP^{-/-}
532 neurons when compared to APP^{+/+} (Extended Data Fig. 5-1), pointing toward a specific
533 impairment in GABA-dependent signaling in the absence of APP. GABA is synthesized in
534 inhibitory neurons by the glutamate decarboxylase enzymes (GAD₆₅ and GAD₆₇). GAD₆₅ is
535 active at nerve terminals and synapses. We observed that GAD₆₅ signal is increased in APP^{-/-}
536 neurons when compared to APP^{+/+} (Fig. 5C). This is not caused by an increase in the relative
537 number of GAD₆₅ positive neurons in APP^{-/-} cultures (Extended Data Fig. 5-2), but likely to an
538 increase in GAD65 expression in GABAergic neurons. To further address the effect of APP
539 deficiency on GABAergic synaptic markers, we first quantified the expression of GABAR α 1, a
540 GABA_A receptor subunit predominantly expressed during neuronal development. We found a
541 slight but significant decrease in GABAR α 1 in APP^{-/-} neurons (Fig. 5D). To evaluate if the
542 activity of GABA receptors was defective in physiological conditions, cortical neurons at DIV7

543 were voltage-clamped at -60mV. To prevent neuronal activity, experiments were performed in the
 544 presence of 1 μ M TTX, 10 μ M CNQX, 20 μ M D-AP5 and 100 nM CGP55845 to inhibit Na⁺
 545 voltage-dependent channels, AMPA, NMDA and GABA_B receptors, respectively. Current-voltage
 546 (IV) curves were generated by evoking the current with a voltage ramp stimulus from -90mV to
 547 +60mV, and the response to 100 μ M was studied (Fig. 5E). Whole cell currents evoked at -50mV
 548 or +50mV by 100 μ M GABA were similar in APP+/+ as in APP-/- neurons (Fig. 5F). This patch-
 549 clamp investigation did thus not reveal any functional decrease in GABA_A receptors, suggesting
 550 that the minor decrease in expression of GABAR α 1 subunit was compensated.

551 Finally, we evaluated the impairment in GABAergic markers in the brain of APP-/- mice. We
 552 quantified the expression of GAD65 and GABAR α 1 in cortices and hippocampi of 3-month-old
 553 mice. A significant increase in GAD65 expression was observed both in cortex (Figure 6A) and
 554 in hippocampus (Figure 6B) of APP-/- mice, and decreased GABAR α 1 levels were measured
 555 especially in the hippocampus. We analyzed CA3-to-CA1 synapse plasticity by extracellular
 556 recordings on hippocampal slices from adult APP+/+ and APP-/- mice. Schaffer collaterals (SC)
 557 were stimulated and fEPSP were recorded in the stratum radiatum of CA1 region. We observed
 558 that the relationship between the stimulus intensity and the fEPSP slope was similar in slices
 559 from both genotypes (Extended Data Fig. 6-1A). To investigate LTP, we stimulated the SC
 560 pathway with a theta- burst stimulation (TBS) consisting of four bursts of five pulses (given at
 561 100 Hz) separated by 200 ms. In slices from APP+/+ animals, TBS induced a large increase of
 562 the fEPSP response size that decayed over the first 20 minutes to a plateau level persisting up
 563 to the end of the experiment. In APP-/- mice, LTP was significantly reduced (Fig. 6C). Typically,
 564 60 min after the TBS, LTP was reduced by half. The response to TBS itself was actually
 565 modified: the second, third and fourth bursts of stimulation were globally decreased in APP-/-
 566 compared to APP+/+, and within each of the four bursts of 5 pulses, the responses to the third,
 567 fourth and fifth stimuli decreased more in slices from APP-/- mice than in APP+/+ (Fig. 6D).

568

569

570 **Silencing NPAS4 mimics APP deficiency in neurons**

571 We used the CRISPR-Cas9 approach in order to directly silence *NPAS4* expression in neurons
572 and analyze whether NPAS4 deficiency could recapitulate a major trait observed in APP-/-
573 neurons, i.e. the upregulation of GABA release and modification of GABAergic markers. The
574 efficiency of CRISPR-Cas9 editing is hard to evaluate by quantifying the mRNA levels of the
575 target gene, since decrease in mRNA could only occur if nucleotide insertions by non-
576 homologous end-joining results in nonsense-mediated RNA decay. Commercially available
577 antibodies poorly detect NPAS4 in basal conditions, but we could still observe that NPAS4
578 protein was diminished upon silencing (CRISPR-*NPAS4* condition, Fig. 7A). We further decided
579 to check the down-regulation of *NPAS4* gene expression by measuring NPAS4 protein upon
580 depolarization by KCl (Lin et al., 2008). In that condition, we found that CRISPR-Cas9-induced
581 silencing resulted in a decrease in NPAS4 by approximately 50% (Fig. 7A). This is comparable
582 to the decrease in NPAS4 mRNAs measured in APP-/- neurons at DIV7 (Fig. 1D). As for
583 CRISPR-Cas9 lentiviral vectors targeting APP, we did not observe toxic effects related to viral
584 transduction of primary neurons (See Extended Data Fig. 3-1C). Strikingly, like APP-deficient
585 primary neurons (Fig. 5C) or brains of APP-/- mice (Fig.6), NPAS4-deficient neurons showed an
586 increase in GAD65 staining (Fig. 7B-C) and GAD65 protein expression (Fig. 7D). Accordingly, a
587 2.5-fold increase in GABA concentration was measured in the medium of primary neurons
588 infected with CRISPR-Cas9 *NPAS4* lentiviruses (Fig. 7C). The expression of GABA receptor
589 subunit alpha 1 (GABARa1) was decreased after *NPAS4* knockdown (Fig.7D), to the same
590 extent as the decrease observed in in APP-/- primary neurons (Fig. 5D).

591

592 Discussion

593 One major function of APP is to control synaptic formation, transmission and plasticity (Muller et
594 al., 2017). We showed here that APP deficiency in cortical neurons impairs the balance between
595 excitatory and inhibitory synaptic markers, and increases GABAergic transmission. This process
596 relies on the activity-dependent transcription factor NPAS4. We initially identified the NPAS4 IEG
597 as downstream APP target by a non-biased transcriptome analysis. The APP-dependent
598 transcription of NPAS4 involves its extracellular domain (sAPP α) and is activated by AICD. APP
599 appears thus to exert a fine tuning of inhibitory synapses in neuronal network. Its absence
600 enhances, through the downregulation of NPAS4, the production and the release of GABA.

601 APP-dependent expression of NPAS4 in differentiated neurons

602 The transcriptome analysis of APP^{+/+} vs. APP^{-/-} neurons at embryonic day 18 (E18-DIV0) and
603 at different stages of primary cortical neuron differentiation (DIV3-DIV7) indicated that the
604 transcriptional changes in the absence of APP were moderate. This unexpected result is in line
605 with a comparative transcriptome study of APP family members in the adult mouse cortex (Aydin
606 et al., 2011). Subtle effects of APP deficiency on the transcriptome could be due to functional
607 redundancies with other APLPs that display similar functions and signaling properties (Shariati
608 and De Strooper, 2013). We did not measure any changes in APLP1 and APLP2 expression in
609 our APP^{-/-} models, in agreement with studies done on total brain extracts (Zheng et al., 1995) or
610 in primary cortical neurons (White et al., 1998). Transcriptional modifications we measured are
611 thus related to the absence of the APP protein *per se*. APP-dependent transcriptional
612 regulations likely act by fine-tuning the expression of classes of gene involved in neuronal
613 pathways rather than robustly regulating single target genes. We found that the expression of
614 the NPAS4 IEG is downregulated in the absence of APP. This particularly at DIV7, when primary
615 neurons start to form a functional network. NPAS4 downregulation was observed in APP^{-/-}
616 primary neurons and upon acute APP knockdown by a CRISPR-Cas9 approach, establishing a

link between APP and NPAS4 transcription. Regarding the possible molecular mechanisms involved here in APP-dependent gene transcription, we found that NPAS4 expression is activated in neurons by AICD expression. It correlates with the fact that DIV7 corresponds to the differentiation stage where AICD is peaking in primary cortical neurons. (Kimberly et al., 2005). However, previous studies showed the secreted ectodomain (sAPP α) is sufficient to rescue functional defects in APP KO mice (Ring et al., 2007; Weyer et al., 2014). We found that treatment of APP $^{+/+}$ neurons (and not APP $^{-/-}$ neurons) with sAPP α significantly increased NPAS4 mRNAs. This observation indicates that (i) the effects of sAPP α require the presence of endogenous APP (ii) homophilic ectodomain interactions are likely to be involved. Soluble APP was suggested to promote its physiological effects by interaction with APP holoprotein (Milosch et al., 2014; Deyts et al., 2016). It is tempting to suggest here that interaction of soluble APP with APP holoprotein present at the cell surface could induce the release of transcriptionally active AICD, but this hypothesis awaits further experimental evidence.

Alteration of GABA release and GABA markers in APP deficient neurons

In the absence of APP, we observed an increase in neuronal outgrowth and GAD65 signal, as well as increased GABA release in the medium of primary neurons. The increase in GAD65 signal was related to an increased signal in GAD65-positive neurons, not to an increased number of GAD65 neurons in our primary cultures. It would be useful to further investigate as to whether an increase in the numbers and functionality of GABAergic neurons occurs in the brain of APP $^{-/-}$ mice. To note, APP was reported to control neurogenesis in adult mice brain (Wang et al., 2014), as process that could account for the modification of neuronal subtypes observed in the absence of APP.

The increase-in GAD65 we observed was in line with increased levels of GABA in the culture medium. *In vivo* studies evidenced increased GABA levels in the brain of APP $^{-/-}$ mice (Lee et al., 2010). However, this elevation of GABA markers was concomitant to a down-regulation in the

642 GABAR α 1 receptor subunit. Recent study also reported that GABAR α 1 is particularly decreased
 643 in hippocampus of APP-/- mice (Chen et al., 2017), in line with our in vivo experiments. Still,
 644 patch-clamp experiments did not reveal any decrease in GABA $_A$ receptors functionality,
 645 indicating that compensation may occur to circumvent the decrease in a GABA $_A$ receptor
 646 subunit, and that GABAergic transmission per se is not significantly altered.

647 On the other hand, our results show that LTP is reduced in brain slices from APP-/- mice. This is
 648 in agreement with previous studies showing that the LTP induced by one TBS is reduced in
 649 APP-/- at the SC-CA1 synapses (Dawson et al., 1999;Seabrook et al., 1999;Ring et al., 2007)
 650 but not at the perforant path-granule cell (DG) synapse that we did not study here (Jedlicka et
 651 al., 2012), Such inhibition was however not observed by other investigators (Wang et al., 2017),
 652 possibly because they used a stronger stimulation protocol (3 TBS). Interestingly, the analysis of
 653 the responses to the TBS itself is consistent with increased GABAergic release in APP-/- mice.

654 How could APP modulate GABA release? Soluble APP (sAPP α) is known to enhance LTP and it
 655 is sufficient to rescue the decrease of LTP observed in APP-/- mice (Taylor et al., 2008;Ring et
 656 al., 2007). Moreover, very recent studies showed that the sAPP α directly binds the GABA $_B$
 657 receptor subunit 1a (GABABR1a), suppressing synaptic transmission and triggering short-term
 658 facilitation in hippocampal neurons (Rice et al., 2019). Such an effect of sAPP α could explain the
 659 smaller response observed in brain slices from APP-/- mice compared to APP+/+. Indeed, in
 660 APP-/- slices, a larger release of GABA would occur in the absence of GABA $_B$ R1a stimulation by
 661 sAPP α , therefore reducing the disinhibitory process observed between the first and the second
 662 burst of stimuli of the TBS (Larson and Munkacsy, 2015). This supports that APP synaptic
 663 function and APP-dependent synaptic transmission are mediated by the soluble sAPP α
 664 fragment. We suggest that the NPAS4 IEG is sensing the APP-dependent modulations of
 665 synaptic transmission. Our results also indicate that NPAS4 knockdown mimics APP deficiency
 666 on GAD65 levels and GABA measurements. The finding that APP functions in neuronal network

might be mediated by NPAS4 is relevant to several evidences about the reported role of NPAS4 in neuronal network activity. NPAS4 possesses unique features among the IEGs (Sun and Lin, 2016): (i) it is only expressed in neurons; (ii) it is activated selectively by neuronal activity; (iii) it has been shown to shape glutamatergic and GABAergic synaptic inputs. NPAS4 is implicated in a transcriptional program that regulates neuronal firing responses to excitatory transmission by enhancing inhibition (Lin et al., 2008), and is critical to keep neuronal firing in response to stimuli within normal levels (Spiegel et al., 2014). This exciting new field of investigation connecting APP function to ITFs sensing neuronal activity awaits further investigation.

Possible relevance to the AD pathophysiology

APP plays a central role in the onset and progression of the amyloid pathology found in AD. Apart from producing A β , the precise contribution of the APP protein to the pathology is poorly understood. The impairment of APP function, either caused by Familial AD (FAD) mutations or upon neuronal ageing, may contribute to neuronal dysfunctions occurring in the disease. In the mammalian brain, APP modulates dendritic complexity, synaptic functions and synaptic plasticity (Muller et al., 2017). A reduction in dendritic length and branching as well as in total spine density was reported in old APP-deficient mice (Lee et al., 2010; Tyan et al., 2012). Aging is an important parameter related to APP functions in the brain. The phenotype related to APP deletion in the central nervous system (CNS) is age-dependent (Priller et al., 2006). Upon aging, impairments in learning and memory associated with deficits in LTP are observed in APP-deficient mice as shown here and in previous studies (Ring et al., 2007). Interestingly, the decline in memory performance and reduction in LTP observed in old mice and APP transgenic mice are mediated by the ionotropic GABA_A receptor (Yoshiike et al., 2008). This imbalance in neuronal excitatory/inhibitory transmission was also observed in the temporal cortex of AD patients, where significantly lower levels of GABA and glutamate were measured (Gueli and Taibi, 2013). These observations unambiguously indicate that changes in neurotransmission

692 occur in AD (and even in ageing brain) and point toward alteration of the inhibitory GABAergic
693 transmission. Important points must be kept in mind here. First, GABAergic transmission shifts
694 from excitatory to inhibitory during development (Ben-Ari, 2002), so the consequence of altered
695 GABAergic transmission can be fully understood only in adult brain. Then, the molecular
696 mechanisms underlying changes in inhibitory transmission in AD are complex. The GABAergic
697 molecular actors are differentially affected by ageing (Vela et al., 2003) or in AD mice models
698 (Yoshiike et al., 2008). Decrease in GABAR α 1 has been reported in ageing rodent brain and in
699 the hippocampus of aged brains with AD (Mizukami et al., 1998), but this could be functionally
700 compensated as shown herein. Our findings should further be addressed in AD mice models
701 (expressing APP mutations) to complete the results obtained here in a loss-of-function model
702 (APPKO). However, the hypothesis of an overall impairment of GABAergic transmission in AD is
703 also supported by the increased risk for unprovoked seizures observed in individuals with AD
704 compared to non-demented individuals of the same age (Friedman et al., 2012).

705 Finally, NPAS4 expression was found to decrease along with AD progression, particularly at
706 Braak NFT stages (I-II) corresponding to lesions developed in transentorhinal/entorhinal cortex
707 (Miyashita et al., 2014). We believe that our main observation, namely that APP deficiency in
708 neurons is integrated by the activity-dependent NPAS4 IEG and affects the balance of inhibitory
709 and excitatory neuronal inputs, provides new insight to understand the role of APP in synaptic
710 activity, but also a mechanistic frame to further explore the impairments of network activity in
711 AD.

712

713 Figure Legends

714 **Figure 1: APP-dependent expression of NPAS4 in young differentiating neuronal culture**

715 Summary of transcriptome analysis performed with the *GeneChip® Mouse Transcriptome Array*
716 *1.0* (Affymetrix). The characterization of the model and the experimental workflow are described
717 in Extended Data Fig. 1-1. Data were processed in triplicate (N=3) for each experimental time
718 point (E18; DIV3; DIV7). Non-coding transcripts and alternative splicing products were detected
719 by these arrays, but only transcripts of coding transcripts have been considered here. For all the
720 transcripts, adjusted p-value > 0.05 except for APP (internal control, p-value < 0.05). **A)** Number
721 of up-and down-regulated coding transcript in APP-/- vs. APP+/+ primary neurons at E18, DIV3
722 and DIV7. Linear fold changes have been set at 1.25, 1.5 and 2. **B)** KEGG pathway analysis
723 (<http://www.genome.jp/kegg/pathway.html>) at E18, DIV3, DIV7 (APP-/- vs. APP+/+) to identify
724 networks molecular pathways (or interaction networks) in which differentially expressed genes
725 are clustered. The five most modified pathways are displayed for each time point, with the
726 number of genes potentially up-or down-regulated. **C)** Immediate Early Genes (IEGs) expression
727 in APP-/- vs. APP+/+ primary neurons at DIV7 and their respective fold change (APP-/- vs
728 APP+/+) in microarray analysis at DIV7. **D)** Neuronal PAS 4 domain (*NPAS4*) mRNA level was
729 measured by qPCR at E18, DIV3 and DIV7 (n=6, N=3). Results (mean ± SEM) are expressed as
730 percentage of controls (APP+/+). n.s.= non-significant, *p=0.0242, Student's t-test. mRNA levels
731 of two other IEGs (*Egr1* and *Egr3*) were measured in the same conditions (Extended Data Fig.
732 1-2).

733 **Figure 2: APP metabolites regulate NPAS4 expression**

734 **A)** Schematic representation of APP, its fragments, the AICD-HA construct along with and the
735 localization of the epitopes recognized by the different antibodies used. Western blotting
736 analysis of AICD-HA expression after 3 days of lentiviral infection in cells with control or AICD-

HA expressing vectors. Total cell lysate was analyzed with anti-HA antibody. **B)** Quantification by qPCR of NPAS4 mRNA in neurons APP^{-/-} at DIV7 infected with lentiviral vector expressing AICD-HA (n=6, N=3). Results are expressed as percentage of control (Ct) (mean± s.e.m). *p.= 0.0291, Student-t test. **C)** Medium of sAPP α treated APP^{+/+} or APP^{-/-} neurons was subjected to Western blotting analysis using anti-human APP antibody (clone W0-2) to detect the exogenous human sAPP α (h sAPP α) and an anti-mouse APP antibody (clone 22C11) to detect both endogenous and exogenous sAPP α (h+m sAPP α). Medium was collected after 16 h of treatment. **D)** Quantification by qPCR of NPAS4 mRNA level in APP^{+/+} (n=8, N=4) or in APP^{-/-} neurons at DIV7 treated with 20 nM sAPP α for 16 h (n=6, N=3). Results are expressed as percentage of control (Ct) (mean ± s.e.m.). **p=0.0055, n.s.= non-significant, Student-t test. Given that primary cultures of cortical neurons at DIV 7 also contain astrocytes, the astrocytic pattern of NPAS4 expression is described in Extended Data Fig. 2-1.

Figure 3: Decreased NPAS4 expression in APP-silenced primary neurons

APP was knockdown by CRISPR-Cas9 approach in primary neurons cultures. The infectivity and toxicity of lentiviral CRISPR-Cas9 vectors are detailed in Extended Data Fig. 3-1. **A)** APP expression characterization in neurons by immunostaining. Cortical neurons were infected at DIV1 with lentiviruses expressing sgRNAs (Oligo2, Oligo17) or not (Ct). All lentiviruses harbor a GFP expression cassette. Cultures were immunostained for MAP2 (red), APP (blue) and DAPI (light blue) at DIV7. Arrowheads indicate the position of GFP-positive (infected) neurons in each condition. Scale bar: 100 μ m. **B)** Quantification of APP signal in GFP-positive neurons by Image-J. At least 33 neurons were quantified in two independent experiments for each condition (n=33 N=2). Results (mean ± SEM) are given as percentage of control (Ct). ###p<0,001 (Ct vs Oligo 2) ***p<0.001 (Ct vs Oligo17); Kruskal-Wallis test and Dunn's multiple comparison test. **C)** Upper panel. Representative Western blots showing APP, APLP1, APLP2 and GAPDH protein levels in cortical neurons at DIV7 infected in the same conditions as in Fig 1A. NI = non-infected. Lower

762 panel. Quantification of APP expression in total cell lysates measured by Western blotting.
763 Results (mean \pm SEM) are given as percentage of control (Ct). *** $p < 0.001$ (Ct vs Oligo17),
764 ### $p < 0.001$ (Ct vs Oligo 2), ANOVA and Bonferroni's multiple comparison test ($n=6$, $N=3$). **D)**
765 Sorting of GFP-expressing neurons (FACS). Scatter plots (FSC vs. SSC, left panels) of non-
766 infected and GFP-expressing cells are shown. Dot plots (TOPRO-3, far red vs. GFP, right
767 panels) were used to gate (green rectangle) GFP-positive/TOPRO-3 negative cells. RNA was
768 extracted from these cells and NPAS4 mRNA level was quantified by qPCR. Results were
769 obtained from pooled samples (4 wells of 4 cm² each) for each condition (Ct, Oligo2 and
770 Oligo17). Quantification was carried out on 2 independent experiments ($N=2$). Results (mean \pm
771 SEM) are expressed as percentage of Ct.

772 **Figure 4: Altered neurites arborization of APP deficient neurons during in vitro maturation**

773 **A)** Cortical APP^{+/+} or APP^{-/-} were stained against the neuron-specific marker MAP2 and the
774 nuclear dye DAPI at different stages of maturation (DIV1-2-3 and DIV7). Scale bar: 400 μ m. **B)**
775 Quantification by ImageJ of MAP2 signal area normalized to the number of neurons at DIV1, 2, 3
776 and 7. Quantifications were from 3 fields of at least 6 coverslips from APP^{+/+} and APP^{-/-}
777 neurons, in three independent experiments ($N=3$). Results (mean \pm SEM) are expressed as
778 percentage of control (APP^{+/+}). * $p=0.0293$, Mann-Whitney test.

779 **Figure 5: GABAergic markers and GABAergic transmission in APP knockout neurons**

780 **A)** Quantification of γ -amino butyric acid (GABA) in culture medium and cell extracts of APP^{+/+}
781 and APP^{-/-} primary neurons at DIV7. Results (mean \pm SEM) are expressed as percentage of
782 APP^{+/+} ($n=20$, $N=3$). ** $p=0.0024$, n.s.= non-significant, Student-t test. **B)** Quantification of
783 glutamate in culture medium and cell extracts of APP^{+/+} and APP^{-/-} neurons at DIV7. Results
784 (mean \pm SEM) are expressed as percentage of APP^{+/+} ($n=16$, $N=3$). n.s.= non-significant,
785 Student-t test. Glutamate responses measured in APP^{-/-} neurons are shown in Extended Data

Fig. 5-1. **C)** Cortical APP^{+/+} and APP^{-/-} neurons at DIV7 were immunostained for the neuron-specific marker MAP2 and glutamate decarboxylase 65 (GAD65). Arrows indicate MAP2/GAD65 positive neurons, shown at higher magnification (insets). The expression profile of the GAD65 GABAergic marker in APP^{+/+} and APP^{-/-} neurons is detailed in Extended Data Fig. 5-2. GAD65 signal (5 fields per coverslip) was normalized for quantification to the number of cells in the area (histogram on the right). At least 2 coverslips were quantified for each group (APP^{+/+} and APP^{-/-}) in two independent experiments (N=2). Results (mean \pm SEM) are given as percentage of control (APP^{+/+}). Scale bar: 200 μ m. *p=0.0220, Mann-Whitney test. **D)** Neurons harvested at DIV7 and cell extracts analyzed by Western blotting for GABAR α 1 and GAPDH expression. Quantification of GABAR α 1 was normalized to GAPDH expression. Results (mean \pm SEM) are expressed as percentage of Ct (n=5, N=2). *p=0.0197, Student's t-test. **E)** Representative I-V traces (from -90 to +60 mV repeated every 0.1 s) through APP^{+/+} (A) and APP^{-/-} (B) neurons, in the presence 100 μ M GABA (red traces). **F)** Pooled data of whole-cell current (at +50 and -50 mV) evoked by 100 mM GABA, through APP WT and KO neurons. Each column represents mean \pm SEM of n=8 cells.

Figure 6: GABAergic markers and LTP in adult mice

A) Left panel: Western blot analysis of GAD65, GABAR α 1 and GAPDH expression in cortex of 3 month old APP^{+/+} and APP^{-/-} mice (N=5). **Right panel:** Quantification of GAD65 and GABAR α 1 were normalized to GAPDH expression. Results (mean \pm SEM) are expressed as percentage of APP^{+/+} (N=5). *p=0.0166, Student's t-test. **B) Left panel:** Western blot analysis of GAD65, GABAR α 1 and GAPDH expression in hippocampus of 3 month old APP^{+/+} and APP^{-/-} mice (N=5). **Right panel:** Quantification of GAD65 and GABAR α 1 were normalized to GAPDH expression. Results (mean \pm SEM) are expressed as percentage of APP^{+/+} (N=5). *p=0.0404, Student's t-test. LTP was measured in hippocampal SC-CA1 pathway from APP^{+/+} (n=9) and APP^{-/-} mice (n=8). The input-output relationship between fEPSP measured in CA1 stratum

811 radiatum and the intensity of SC stimulation is depicted in Extended Data Fig. 6-1. **C)** fEPSP
812 slopes measured during TBS ($p<0.05$; two-way repeated measurements ANOVA, Bonferroni).
813 **D)** fEPSP slopes measured before and after TBS. Results expressed in proportion of the
814 baseline response (100%). ($p<0.05$; two way repeated measurements ANOVA, Bonferroni).

815 **Figure 7: NPAS4 silencing by CRISPR-Cas9 mimics cell phenotype observed in APP**
816 **deficient neurons.**

817 Changes on inhibitory (GABA) synapses was analyzed after *NPAS4* silencing **A) Left panel.**
818 Cortical neurons infected with CRISPR-Cas9 lentivirus targeting *NPAS4* gene (CRISPR-*NPAS4*)
819 show reduced *NPAS4* levels as measured by Western blotting (high exposure). Same
820 experiments were carried out after membrane depolarization with 50mM potassium chloride
821 (KCl). *NPAS4* accumulations was detectable by Western blotting at low exposure. Viruses
822 without sgRNA were used as controls (Ct). **Right panel.** Quantification of *NPAS4* protein level
823 after 2, 3 and 4h of KCl depolarization. Results (mean \pm SEM) are expressed as percentage of
824 non-treated controls Ct (N=2). *** $p<0.0001$ Student's t-test. **B)** Cortical neurons infected with
825 CRISPR-*NPAS4* lentiviruses at DIV1 were immunostained against MAP2 and glutamate
826 decarboxylase 65 (GAD65) at DIV7. Quantification of GAD65 signal was normalized to the
827 number of cells (5 fields per coverslip, 2 coverslips for each genotype in two independent
828 experiments, (N=2). Results (mean \pm SEM) are given as percentage of control (Ct). Scale bar:
829 200 μ m. ** $p=0.0024$. Mann-Whitney test. **C)** Quantification of γ -amino butyric acid (GABA) in
830 culture medium at DIV7 of infected control neurons (Ct) and CRISPR-*NPAS4* infected neurons.
831 Results (mean \pm SEM) are expressed as percentage of Ct (n=5, N=2). * $p=0.0146$, Student-t test.
832 **D)** Neurons harvested at DIV7 and cell extracts analyzed by Western blotting for GABAR α 1,
833 GAD65 and GAPDH expression. Quantification of GABAR α 1 and GAD65 were normalized to
834 GAPDH expression. Results (mean \pm SEM) are expressed as percentage of Ct (n=8, N=3).
835 * $p=0.049$, # $p=0.0247$, Student's t-test.

836

837 **Extended Data Figure Legends**838 **Figure 1-1: Experimental workflow and model characterization**

839 **A)** Experimental design used for the study (E18); neurons were cultured and experiments were
840 mainly carried out after 3 and 7 days in vitro (DIV3 and DIV7). Transcriptome analysis was
841 performed on embryonic cortex (E18) and at DIV3 or DIV7. **B)** APP, APLP1 and APLP2
842 expressions were analyzed by Western blotting at the indicated days of culture in APP+/+
843 neurons. **C)** Quantification of APP, APLP1 and APLP2 protein expression over time in APP+/+
844 neurons. Accumulation is represented as fold change over the signal measured at day 0.
845 Quantification was performed from one neuronal culture **D)** APLP1 and APLP2 expressions are
846 not modified in cortical tissue at E18 and primary neuron cultures at DIV3 and DIV 7 in absence
847 of APP. Expression of APP, APLP1, APLP2 was analyzed by Western blotting of cells lysates
848 from APP+/+ and APP-/- primary neuron cultures. **E)** Samples from primary cultures at DIV7
849 (APP+/+, APP+/- and APP-/- neurons) were probed (Western blotting) with an antibody directed
850 against APP C-terminus for APP C-terminal fragments (CTFs) and AICD. Low and high
851 exposures of a typical blot are shown. Arrows indicate the expected position of APP holoprotein,
852 APP CTFs and AICD.

853 **Figure 1-2: Expression of Egr1 and Egr3 is not modified in APP-/- neurons**

854 Egr1 and Egr3 expressions were evaluated in APP+/+ vs. APP-/- primary neurons at DIV7. **A)**
855 Egr1 mRNA and **B)** Egr3 mRNA levels was measured by qPCR (n=6, N=3) at DIV7. Results
856 (mean \pm SEM) are given as percentage of controls (APP+/+) n.s= non-significant, Student's t-
857 test.

858

859 **Figure 2-1: Astrocytes in primary neuron culture and their implication in Npas4**
860 **expression.**

861 **A)** Primary culture of cortical neurons at DIV7. Cultures were immunostained with the neuron
862 specific protein MAP2 (green), the glial specific protein GFAP (red) and the DAPI (light blue).
863 Scale bar = 400µm. **B)** Quantification of neurons (MAP2+) and astrocytes (GFAP+) in the
864 primary cortical culture. At least five fields per coverslip were analyzed for APP+/+ and APP-/-
865 cultures in two independent experiments (n≥5, N=2). Results are expressed as the ratio of
866 MAP2+ (neurons) and GFAP+ (astrocytes) (mean ± s.e.m). n.s= non-significant, Mann-Whitney
867 test. **C)** Western blotting analysis of NPAS4 induction in neurons and astrocytes after
868 depolarization with 50mM potassium chloride (KCl) for 2 hours.

869 **Figure 3-1: Infectivity and toxicity of lentiviral CRISPR-Cas9 vectors**

870 **A)** Cortical neurons were infected at DIV1 with lentiviruses expressing sgRNAs (Oligo2, Oligo17
871 or CRISPR-NPAS4) or no sgRNA (Ct), SpCas9 and GFP. Cultures were immunostained for
872 MAP2 (red) and DAPI (light blue) at DIV7. Scale bar = 400µm. **B)** Quantification of GFP+
873 neurons (GFP+/MAP2+) in total neuron population (MAP2+) after lentiviral CRISPR-Cas9
874 infection with control (Ct), Oligo2, Oligo17 or CRISPR-NPAS4. At least five fields were analyzed
875 for each lentiviral vector in two independent experiments (n≥5, N=2). Results are expressed as
876 percentage of GFP+/MAP2+ cells in total MAP2+ cells (mean ± s.e.m). n.s= non-significant,
877 Kruskal-Wallis test and Dunn's multiple comparison test. **C)** Measurement of LDH activity
878 released after infection (DIV7) of primary neuron with control (Ct), Oligo2, Oligo17 or CRISPR-
879 NPAS4 at DIV7 lentiviral vectors. Background LDH release was determined in non-infected
880 control cultures (NI). Results were expressed as percentage of total LDH release measured in
881 non-infected control cultures (NI) in 2 independent experiments (n=12, N=2).

882 **Figure 5-1: Glutamate responses in APP-/- neurons measured by calcium imaging.**

Neuronal activity was measured at DIV7 by calcium imaging. **A)** Left panel. Different calcium responses were observed after stimulation with 50 μ M glutamate and classified as described by Prickering and co-workers (Prickering et al. 2008) between neuronal and non-neuronal responses. To note X-axis graduation correspond to 20 sec. Right panel. The proportion of cells displaying Type 1, 2 or 3 response was quantified in three independent experiments (n=9, N=3). n.s.= non-significant. Student-t test. **B)** Normalized fluorescence trace (mean \pm SEM) measured in APP^{+/+} and APP^{-/-} neurons upon perfusion for 150 sec with 50 μ M glutamate. The area under curve (AUC) was quantified for 50 neurons per coverslips. A total of 9 coverslips for each genotype was recorded in three independent experiments (N=3). The graph on the right shows AUC expressed as percentage of control (APP^{+/+}). *p=0.0106, Student's t-test.

Figure 5-2: GAD65 positive neurons in primary cortical cultures.

A) Primary culture of cortical neurons after at DIV7. Cultures were immunostained with the neuron specific protein MAP2 (red), GAD65 (dark blue) and DAPI (light blue). Representative 20x micrographs show GAD65 positive neurons (white arrowhead) and GAD65 negative neuron (green arrowhead). **B)** Images (20x objective) were quantified (10 fields per coverslip for each genotype) in three independent cultures (n=30, N=3). Results (mean \pm s.e.m) are expressed as percentage of GAD65+ MAP2+ cells (GAD65+ neurons) among all MAP2+ cells (neurons). n.s= non-significant, Mann-Whitney test. Scale bar = 20 μ m.

Figure 6-1: LTP in hippocampal SC-CA1 pathway in APP^{-/-} mice

Excitatory postsynaptic potentials measured in hippocampal CA1 region of brain slices from APP^{+/+} (N=9) and APP^{-/-} mice (N=8). **A)** The input-output relationship between fEPSP measured in CA1 stratum radiatum and the intensity of SC stimulation is represented. No significant difference between APP^{+/+} and APP^{-/-} was observed.

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Figure 1

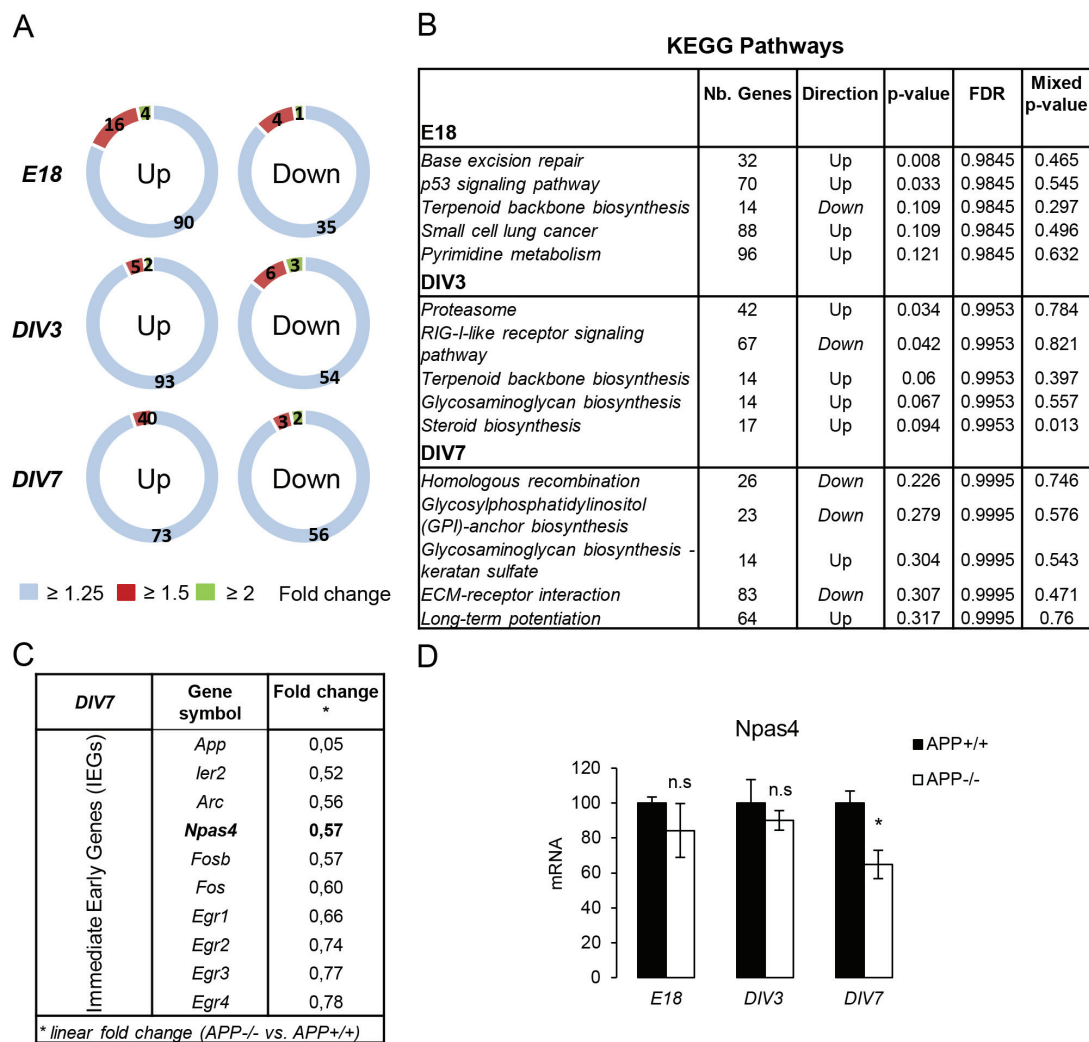


Figure 2

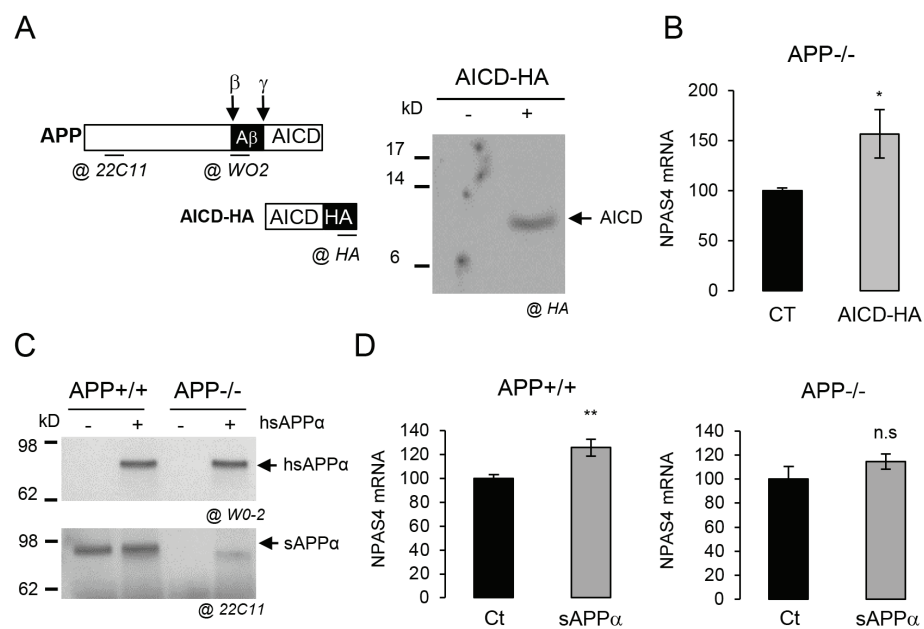


Figure 3

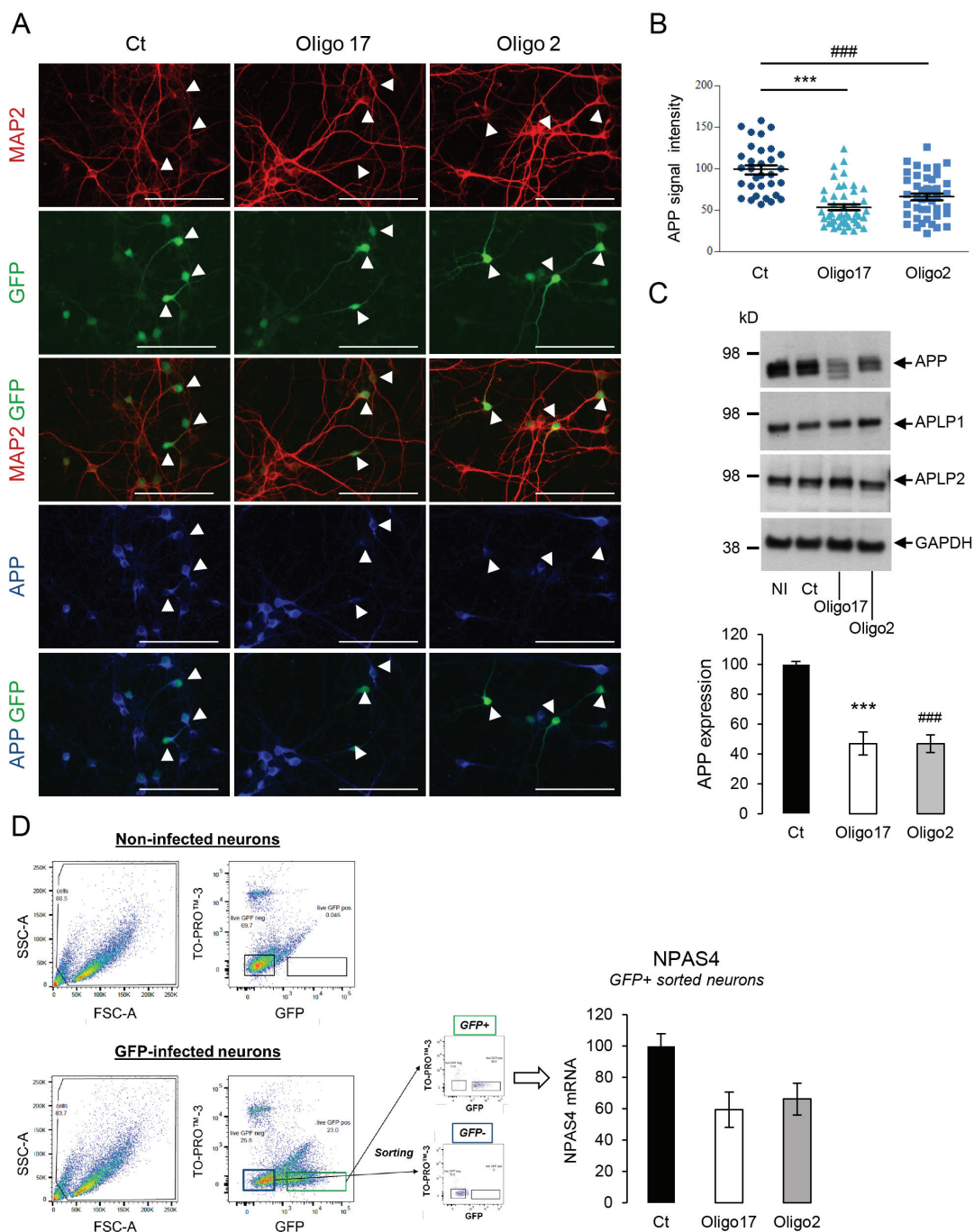


Figure 4

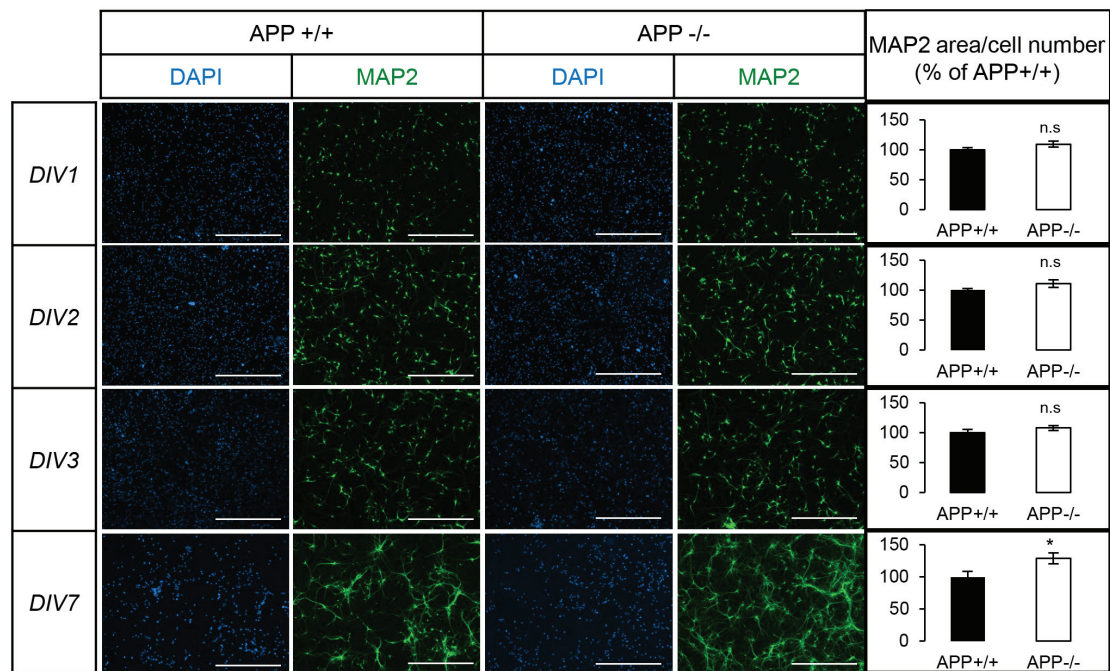


Figure 5

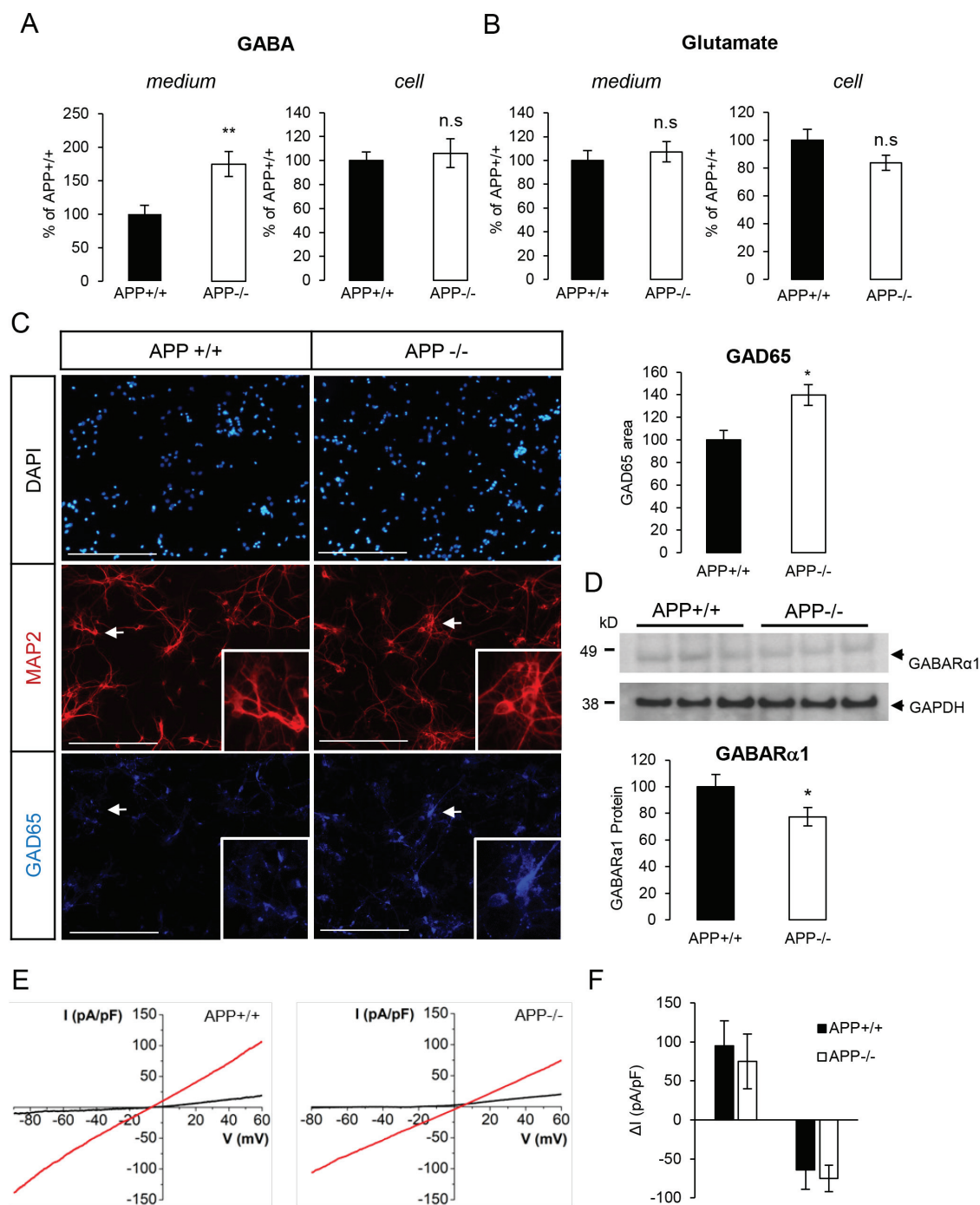


Figure 6

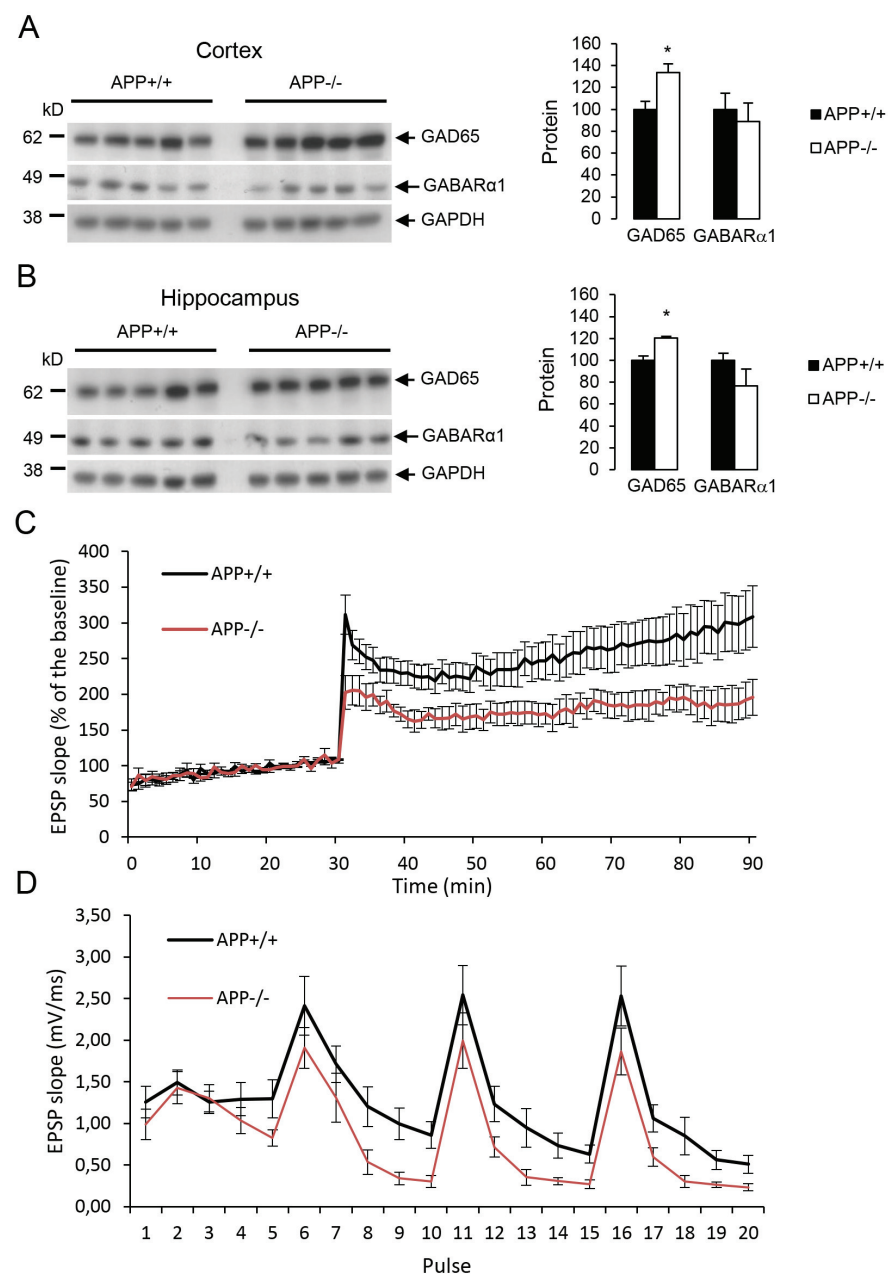


Figure 7

